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(54) Title: COMPOSITIONS AND METUODS	FOR C	7.0	NES CONTAINING DNA SEQUENCES ASSOCIATED

(54) Title: COMPOSITIONS AND METHODS FOR CLONES CONTAINING DNA SEQUENCES ASSOCIATED WITH MULTIDRUG RESISTANCE IN HUMAN CELLS

(57) Abstract

Genomic and cDNA clones of human genes which are selectively amplified or overexpressed in multidrugresistant human tumor cells were isolated. Such clones may be used as probes in diagnostic tests to detect chemotherapy-resistant tumor cells and to predict tumor response to chemotherapy. The complete nucleotide sequence of the coding region of the human mdrl gene and the complete corresponding amino acid sequence are disclosed.

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COMPOSITIONS AND METHODS FOR CLONES CONTAINING DNA SEQUENCES ASSOCIATED WITH MULTIDRUG RESISTANCE IN HUMAN CELLS

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Roninson et al.

This is a Continuation-in-Part of Application Serial No. 845,610, filed March 28, 1986.

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Background

The present invention pertains in—general to diagnostic materials and methods and in particular to materials and methods for the detection of multidrug-resistant tumor cells.

Selection of mammalian cells for resistance to plant alkaloids or antitumor antibiotics frequently results in the development of cross-resistance to other drugs unrelated in their structure and mode of action to the original selective agent. Biedler et al., Cancer Res., 30, 1174 (1970). The phenomenon of multidrug resistance constitutes a major problem in cancer chemotherapy since it involves resistance to some of the most commonly used anticancer drugs.

Multidrug resistance in most cases appears to result from decreased intracellular drug accumulation, probably as a result of alterations in the plasma membrane. Biedler et al., Cancer Treat. Rep., 67, 859 (1983); Ling et al., Cancer Treat. Rep., 67, 869 (1983); 30 Ramu et al., Cancer Treat. Rep., 67, 895 (1983); and Beck et al., Cancer Res., 39, 2070 (1979).

In some hamster, mouse and human multidrugresistant cell lines, resistance correlates with over
expression of a 170,000 m.w. membrane glycoprotein (Pglycoprotein) or a 19,000 m.w. cytosolic protein. Kartner et al., Science, 221, 1285-1288 (1983); Biedler et

al., Cancer Treat. Rep., 67, 859 (1983). Immunoblotting techniques applied to cells from human cancer patients reveal high levels of P-glycoprotein in some cases of advanced, nonresponsive ovarian cancer. Bell et al.,
5 J. Clin. Oncol., 3, 311-315 (1985).

P-glycoprotein-specific, monoclonal antibodies raised against multidrug-resistant Chinese hamster ovary (CHO) cell lines and cross reactive with human cell lines apparently bind to multidrug-resistant mammalian 10 cells to a degree correlated with the degree of their drug resistance. - Kartner et al., Nature, 316, 820-823 (1985). These monoclonals may all bind to a C-terminal intracellular region of a proposed P-glycoprotein polypeptide. Kartner et al., Nature, 316, 820-823 (1985). P-glycoprotein specific cDNA clones have been 15 isolated from Chinese hamster ovary cells, and these clones revealed amplification of the P-glycoprotein gene in multidrug resistant hamster, mouse and human cells when employed in a Southern blotting procedure. Riordan 20 et al., Nature, 316, 817-819 (1985). However, Riordan et al. provides no indication whether the hamster Pglycoprotein cDNA clones may be used to detect the expression of human P-glycoprotein genes at the level of RNA.

In a different approach to the examination of multidrug-resistance, a common region of DNA is found to be amplified in two different multidrug-resistant Chinese hamster cell lines selected for resistance to either colchicine or Adriamycin. Roninson et al.,

Nature, 309, 626 (1984). This region was found to contain a transcription unit, presently designated mdr. Expression of the mdr mRNA correlates with multidrug resistance in the hamster cells. Gros et al., J. Cell. Biochem., 9C (suppl.), 16, All67 (1985); and Gros et al., Proc. Natl. Acad. Sci. (USA), 83, 337 (1986). However, probes derived from the hamster mdr. gene are

not useful probes for human cells inasmuch as, even though these probes hybridize to human DNA (as illustrated in Example 2, <u>infra</u>), they do not hybridize efficiently with human <u>mdr</u> mRNA, despite the impression given in a report on a workshop dealing with multidrug resistance [Kolata, <u>Science</u>, 231, 220-221 (1986)].

Therefore, in the absence of a probe for human mdr gene expression, there is a need for a reliable method for detecting the presence of multidrug-resistant cells in a human tumor either prior to or during chemotherapy.

Summary of the Invention

The present invention provides an isolated nucleic acid sequence for a human mdr gene associated with multidrug resistance in human cells.

A presently-preferred embodiment of the present invention provides an isolated and purified nucleic 20 acid selected from the group consisting of: (a) a nucleic acid comprising a member of the group consisting of a continuous sequence of nucleotides as set forth in Table 4, in Table 5, in pHDR4.4 (ATCC 4Q227), in pHDR4.5 (ATCC 40228), in pHDR5A (ATCC 67040), in pHDR5B (ATCC 25 67041), in pHDR10 (ATCC 67042) and in pHDR104 (ATCC 67156); (b) a nucleic acid comprising a nucleotide sequence which hybridizes with at least one of the continuous sequences of nucleotides as set forth in (a) or which is contained within the same mRNA molecule of 30 human origin or cDNA molecule of human origin as at least one of the continuous sequences of nucleotides as set forth in (a); (c) nucleic acids comprising a nucleotide sequence which hybridizes with any nucleotide sequences described in (b); and (d) nucleic acids comprising a sequence of nucleotides which, but for the degeneracy of the genetic code, would hybridize

with at least one of the continuous sequences of nucleotides as set forth in (a), (b), or (c). Standard conditions for identifying the presence or absence of "hybridization" herein are reactions conducted in 4 X 5 SSC and 0.5% SDS at a temperature of 65 degrees C. in the last wash. A nucleic acid probe according to the preferred embodiment may also include a label associated with one of these nucleic acids. Polypeptides encoded by these nucleic acids may be expressed or synthesized 10 chemically, and used, in conjunction with diluents, adjuvants, or carriers of the sort well known to those skilled in the art, to raise monoclonal or polyclonal antibodies or to elicit immune response in patients. Such antibodies may be utilized as a diagnostic reagent 15 using various presently available immunodiagnostic techniques, or employed as immunotherapeutic agents.

Brief Description of the Drawings

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Fig. 1 is a partial restriction map of the cosmid clone cosDR3A which contains a 5' portion of the transcribed mdr region isolated from Chinese hamster cells:

Fig. 2 illustrates partial restriction maps of the plasmid clones pHDR4.4 and pHDR4.5, respectively containing mdrl and mdr2 sequences; and

Fig. 3 illustrates partial restriction maps of phage cDNA clones λHDR5, λHDR10, λHDR62, λHDR28, 30 λHDR69A, λHDR103 λHDR104 and λHDR105 containing mdr1 sequences.

Detailed Description

35 Preliminary announcements of the obtaining of mdrl clones according to the present invention and of

uses therefor have been made by the inventors at the UCLA Symposia on Molecular and Cellular Biology, January 20 - February 15, 1986. Roninson et al., J. Cell.

Biochem., 29 (suppl. 10A), 12, Al8 (1986); Pastan et al., J. Cell. Biochem., 29 (suppl. 10A), 9, Al3 (1986); Clark et al., J. Cell. Biochem., 29 (suppl. 10A), 49, Al30 (1986); and Cornwell et al., J. Cell. Biochem., 29 (suppl. 10A), 50, Al31 (1986).

A recently published European Patent 10 Application No. 174,180 by John R. Riordan, entitled "Multidrug Resistance in Mammalian Cell Lines And Isolation Of Determinant Glycoprotein DNA," describes isolation of Chinese hamster cDNA clones specific for Pglycoprotein, and it suggests using P-glycoprotein-15 specific cDNA as a probe in determining multidrug resistance in cells. Although only Southern blot hybridization between hamster cDNA and human genomic DNA is described, claim 18 of Riordan, EPA 174,810, relates to a P-glycoprotein-specific DNA molecule "derived from a source selected from the group consisting of Chinese Hamster Ovary cells, mouse cells and human cells." the event that the mdr clones described herein represent the human P-glycoprotein gene sequences, which is likely to be the case as discussed in Example 10 below, it

In fact, Riordan, EPA 174,810, post-dates the publication of Roninson et al., Nature, 309, 626 (1984) which described cloning of a segment of the Chinese hamster mdr region. The work describe in Roninson et al., Nature, 309, was followed by isolation of the entire Chinese hamster mdr gene [Gros et al., J. Cell. Biochem. and Proc. Nat'l. Acad. Sci. (USA), supral as opposed to only partial cDNA clones of the Chinese hamster P-glycoprotein genes, as described in Riordan, EPA 174,810. Riordan, EPA 174,810, provides no evidence

should be noted that Riordan, EPA 174,810, does not disclose a human mdr gene or any portion thereof.

for the ability of Chinese hamster clones to detect the expression of human P-glycoprotein mRNA. Furthermore, the use of P-glycoprotein cDNA as a probe for detection of multidrug resistance in tumor cells is described in 5 Riordan, EPA 174,180, only in terms of detection of amplified P-glycoprotein genes but not in terms of detection of increased P-glycoprotein mRNA expression. Increased mRNA expression, as described in Example 7 below, provides a much more useful diagnostic marker for multidrug resistance than does gene amplification. addition, although claiming P-glycoprotein cDNA sequences of human origin, Riordan, EPA 174,810, contains no indication as to how such sequences would be obtained, e.g. the source of human DNA or RNA, or stringency conditions for screening of human cDNA or genomic libraries with a Chinese hamster probe. As shown in Example 2 below, there is a low level of homology between the hamster and human mdr genes, at least within the 5' half of the gene, which presents a 20 considerable technical problem in the isolation of human mdr DNA sequences.

In the following examples, nucleic acid clones for human mdr genes and uses for the nucleotide sequences of mdr clones are described. In Example 1 a 25 Chinese hamster mdr clone is used to identify sequences hybridizing with human DNA. Example 2 describes the identification and isolation of DNA sequences comprising human mdr genes. In Example 3, amplification of mdr genes in human drug-resistant cells is demonstrated. A 20 characterization of clones containing mdr sequences is presented in Example 4. In Example 5, DNA rearrangement involving mdr genes is examined. In Example 6, transcription of the mdrl gene in human cells is demonstrated. Example 7 describes an investigation into expression levels of the mdrl sequence during the course of development of multidrug resistance in human cells.

In Example 8, expression of mdr genes out of proportion to gene amplification is demonstrated. Example 9 provides a description of a genomic clone containing a segment of the mdrl gene. In Example 10, cDNA clones of the mdrl gene and the cDNA sequence of the human mdrl gene is disclosed are described. In Example 11, diagnostic and therapeutic procedures using probes according to the present invention are described.

10

Example 1

Derivation and characterization of multidrugresistant sublines of human KB cells are described

15 elsewhere. Akiyama et al., Somat. Cell Mol. Genet., 11,
117 (1985); Fojo et al., Cancer Res., 45, 3002 (1985);
and Richert et al., Proc. Natl. Acad. Sci. (USA), 82,
2330 (1985). The multi-drug resistant phenotype is
unstable in the most highly resistant lines, with a

20 decrease in resistance when grown in the absence of the
drugs. Using the in-gel DNA renaturation technique
[according to Roninson, Nucleic Acids Res., 11, 5413
(1983)], several of the multidrug-resistant sublines of
KB cells are known to contain amplified DNA sequences,
25 and karyotypic analysis reveal double minute chromosomes
in these cells. Fojo et al., Proc. Natl. Acad. Sci.
(USA), 82, 7661 (1985).

Sublines of the human KB carcinoma cells, selected for resistance to colchicine, vinblastine or Adriamycin (Akiyama et al., supra; Fojo et al., Cancer Res. supra; Richert et al., supra and Shen et al., Science, 232, 643-645 (1986)], demonstrate the multidrug-resistant phenotype. Several of these sublines are described in Table 1. Fojo et al., Proc. Natl. Acad. Sci. USA, 82, 7661 (1985). In Table 1, "n/d" means not determined. KB-8-5-11, KB-8-5-11-24,

KB-C3 and KB-C4 cell lines are subclones selected in 100 ng/ml, $1\mu g/ml$, $3\mu g/ml$ and $4\mu g/ml$ Adriamycin, respectively. Relative resistance is expressed as the D_{10} of the resistant cell line divided by the D_{10} of the parental KB-3-1 cells. Akiyama et al., supra.

TABLE 1

		Relati	ive Resistance	To:
10	Cell Line	Colchicine	Adriamycin	Vinblastine
	KB-3-1	1		1
	KB-8-5-11	40	23	51
	KB-8-5-11-24	128	26	20
15	KB-C3	487	141	. 206
	KB-C4	1750	254	159
	KB-C1-R1	6	3	4
	KB-V1	171	422	213
	KB-Al	19	97	43
20	KB-A2	n/d	140	n/d

were used to determine whether DNA sequences homologous to the hamster mdr gene are present in the human genome. The Chinese hamster mdr DNA sequences used in this study were derived from the cosmid clone cosDR3A, containing a 5' segment of the hamster mdr gene. After digestion with the restriction enzymes XbaI and KpnI, individual 1.5 - 6 kilobase (kb) restriction fragments from this cosmid were either subcloned into pSP65 plasmid vector commercially available from Promega Biotec, Madison, Wisconsin, or gel-purified prior to labeling with 32p. A vector including a 4.7 kb XbaI fragment, designated pDR4.7, contained DNA sequences hybridizing to human DNA.

In Fig. 1, a partial restriction endonuclease map of the cosmid clone cosDR3A, containing a 5' portion of the transcribed mdr region amplified in multidrugresistant Chinese hamster cells, is presented along with a dashed line aligned to indicate the portion of ROS DR3A which hybridizes to pDR4.7. In Fig. 1, X denotes an XbaI site and K identifies a KpnI site. Cloning and characterization of this region are described in Gros et al., Proc. Natl. Acad. Sci. USA, 83, 337 (1986).

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Example 2

In order to identify and isolate segments of DNA comprising the human mdr genes, individual 1.5 - 6 kilobase (kb) size fragments of the cloned hamster mdr gene were isolated as a series of recombinant subclones in a pSP64 plasmid vector commercially available from Promega Biotec and described in Promega Biotec Technical Bulletin No. 13 as well as in Melton, Nucleic Acids

Res., 12, 7055-7056 (1984). Individual subclones were then labeled with 32p and were used as probes for Southern blot hybridization with human DNA digested with restriction enzymes.

The subclones were then used as probes for hybridization with restriction digests of human genomic DNA. Most probes, when used under conditions of low hybridization stringency, produced either no hybridization signal or a continuous smear suggesting crosshybridization with human repetitive DNA sequences.

30 However, one of the subclones, designated pDR4.7 and illustrated in Fig. 1, gave rise to distinct bands when hybridized to human DNA under low stringency conditions.

Inasmuch as subclone pDR4.7, produced a distinct hybridization signal, this subclone contained hamster DNA sequences homologous to the human mdr genes. pDR4.7 hybridized to two major different EcoRI

5

restriction fragments in human DNA, although in some experiments as many as nine additional EcoRI fragments could be detected.

Example 3

In order to determine whether an mdr gene is amplified in multidrug-resistant human cells, DNA extracted from the parental KB-3-1 cells and various.

10 multidrug-resistant sublines described in Table 1 by the procedure of Gros-Bellard et al., Eur. J. Biochem., 36, 32 (1978) was digested with EcoRI or HindIII, electrophoresed on agarose gels and hybridized to the pDR4.7 probe by the procedure of Southern [Southern, J. Mol. Biol., 98, 503, (1975)].

In the Southern hybridization of pDR4.7 with EcoRI-digested DNA from multidrug-resistant KB cells, DNA was extracted as previously described [Gros-Bellard et al., Eur. J. Biochem., 36, 32 (1978)]. The concen-20 tration of EcoRI-digested DNA was determined by the diphenylamine reaction [Giles et al., Nature, 206, 93 (1965)] and 5 ug of DNA were loaded onto each lane. After electrophoresis, DNA was transferred onto a nylon (Biodyne) membrane [Southern, supra]. Plasmid pDR4.7 25 was digested with XbaI, the insert was gel-purified and labeled with 32 P to a specific activity of 3 x 10⁹ dpm/µg by oligolabeling [Feinberg et al., Analyt. Biochem., 132, 6 (1983)]. Hybridization was done at 65°C in 5 x SSPE, 5 x Denhardt's, 0.2% SDS, 500 μ g/ml 30 denatured salmon sperm DNA. After hybridization, the membranes were washed with 4 x SSC, 0.5% SDS at 65°C and autoradiographed.

The subclone pDR4.7 hybridizes to two <u>EcoRI</u>
fragments of 13.5 and 4.5 kb size and to two <u>HindIII</u>
35 fragments of 10.5 and 4.4 kb size in KB-3-1 DNA when the filters are washed under low stringency conditions (4 x

SSC; 65°C). Only the 13.5 kb EcoRI and 4.4 kb HindIII fragments were detectable under conditions of intermediate stringency (1 x SSC; 65°C). All the fragments were amplified in colchicine-resistant sublines

5 KB-8-5-11, KB-8-5-11-24, KB-C3 and KB-C4.

No amplification of either the band corresponding to the 13.5 kb fragment or the band corresponding to the 4.4 kb fragment was detected in the revertant subline KB-Cl-Rl. Unlike the colchicine-selected sublines, the subline KB-V1, selected in vinblastine, shows amplification of only the 13.5 kb EcoRI and the 4.4 kb HindIII bands. These two bands were also amplified in Adriamycin-resistant cells KB-Al and KB-A2. KB-Al, in addition, contained a new amplified band of a 7 kb size in the EcoRI digest and of a 6.5 kb size in the HindIII digest. The same bands were present in KB-V1 DNA, but their intensity suggested that these bands were not amplified. No bands of this size were detected in the parental KB-3-1 DNA, suggesting that they apparently

20 arose as a result of a DNA rearrangement.

The different patterns of amplification of the two types of bands hybridizing to the hamster mdr probe in different sublines suggested that they might correspond to two different related DNA sequences, possibly different members of a multigene family, rather than to two different parts of one contiguous hybridizing region. DNA sequences corresponding to the 13.5 kb ECORI and 4.4 kb HindIII fragments were designated mdrl and the sequences corresponding to the 4.5 kb ECORI and the 10.5 kb HindIII fragments were designated mdr2.

The degree of amplification of mdr sequences in different multidrug-resistant sublines was estimated by comparing the intensity of hybridization signals from serially diluted EcoRI digests of different cellular DNAs. The estimates of the copy number of mdr sequences in different sublines are given in Table 2. In Table 2,

a star indicates the rearrangement of mdr2 DNA sequences.

TABLE 2

5	Cell Line	Degree of mdrl	Amplification mdr2
	3011		
	KB-3-1	1	1
10	KB-8-5-11	7-8	7-8
	KB-8-5-11-24	9	. 9 .
	квс3	20	20
	KB-C4	30	30
	KB-C1-R1	1	1
15	KB-V1	100	1*
	KB-Al	70	30*
	KB-A2	80	1

By comparison of Table 1 with Table 2, it may 20 be observed that in the sublines selected for a 40-700 fold degree of resistance to colchicine, there is a general, but not precise, correlation between increases in drug resistance and in the copy number of \underline{mdr} sequences. The degree of resistance may correlate more 25 precisely with the expression of mdr RNA than with the degree of mdr gene amplification. The mdrl and mdr2sequences appear to be amplified to a similar degree in these cells. The loss of amplified mdr sequences in a revertant of a colchicine-resistant cell line provides 30 strong additional evidence that mdr gene amplification underlies multidrug resistance in the highly resistant cells. The degree of amplification of mdrl in the cells selected for resistance to vinblastine or Adriamycin appears to be higher than in the cells with a similar degree of resistance that have been selected with colchicine.

XbaI.

Example 4

To investigate the nature of the human mdr 5 genes, clones containing mdrl and mdr2 sequences were isolated from the DNA of the colchicine-resistant subline KB-C3. For this purpose, two phage libraries containing complete EcoRI or HindIII digests of KB-C3 DNA were prepared. The EcoRI library was constructed by 10 insertion into the EcoRI site of the Agtll phage vector, and the HindIII library was made by insertion into the HindIII site of Charon 28 [Young et al., Proc. Natl. Acad. Sci. (USA), 80, 1194 (1983); Rimm et al., Gene, 12, 301 (1980)]. Both libraries were screened by plaque 15 hybridization with the Chinese hamster pDR4.7 probe according to the procedure of Benton et al., Science, 196, 180 (1977). A clone containing the 4.4 kb HindIII fragment (mdrl) was isolated from the HindIII library, and a clone containing the 4.5 kb EcoRI fragment (mdr2) 20 was isolated from the EcoRI library. Both inserts were subsequently recloned into the plasmid vector pSP64 [Melton et al., Nucleic Acids Res., 12, 7035 (1984)], giving rise to plasmid clones designated pHDR4.4 and pHDR4.5, respectively. Plasmid clone pHDR4.4 was 25 deposited with the American Type Culture Collection, 12301 Parklawn Drive, Rockville, Maryland, as Deposit No. 40227 on March 21, 1986. Likewise, plasmid clone pHDR4.5 was deposited with the American Type Culture Collection, 12301 Parklawn Drive, Rockville, Maryland, 30 as Deposit No. 40228 on March 21, 1986. Partial restriction maps of these clones are shown in Fig. 2. In Fig. 2, sites for digestion by corresponding restriction endonucleases are identified as follows: "A", <u>Ava</u>I; "B", <u>Bam</u>HI; "E", <u>Eco</u>RI; "G", <u>Bg1</u>II; "H", 35 HindIII; "J", HaeII; "P", PstI; "V", PvuII; and "X",

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In Fig. 2, solid bars indicate the fragments containing highly repeated sequences. These fragments were identified by hybridization of Southern blots containing restriction digests of cloned DNA with 0.35 x 10⁵ dpm/cm² of ³²P-labeled total human genomic DNA. Dashed lines indicate the DNA sequences hybridizing to the pDR4.7 clone, as determined by Southern hybridization with the gel-purified pDR4.7 insert.

Because the pDR4.7 hamster probe was known to contain transcriptionally active sequences expressed in multidrug-resistant hamster cells (Gros et al., Proc. Nat'l. Acad. Sci. (USA), supra | it seemed likely that the conserved human mdr sequences would provide convenient probes for transcription studies. The 15 hamster pDR4.7 probe hybridized very poorly, if at all, to mRNA from multidrug-resistant human cells, and therefore could not be used as a probe for detection of mdr genes in human cells. Consequently, repeat-free fragments of both clones which hybridized to pDR4.7 were 20 subcloned into the plasmid vector pSP64. The clone containing a 0.75 kb PvuII fragment of pHDR4.4, inserted into the Smal site of the vector, was designated pMDR1. The clone containing a 1.0 kb PstI fragment of pHDR4.5, inserted into the PstI site of the vector, was designated pMDR2. These two clones were found to crosshybridize with each other under conditions of low hybridization stringency providing additional evidence that mdrl and mdr2 represent related DNA sequences.

30

Example 5

To determine whether the rearranged bands in KB-V1 and KB-Al correspond to mdrl or mdr2, DNA from different sublines was digested with HindIII and hybridized to either hamster pDR4.7 probe or to the human

pMDRl or pMDR2 probes. Hybridization with the gel-purified insert of the plasmid pDR4.7 was done under conditions of low stringency (4 X SSC, 0.5% SDS at 65°C). The same blot was then rehybridized with gel-purified inserts of the plasmids pMDRl and pMDR2 under high stringency conditions (0.1 X SSC, 0.5% SDS at 65°C) so that the signal resulting from cross-hybridization of mdrl and mdr2 sequences was minimized.

This experiment demonstrated that rearranged 10 bands in both KB-Al and KB-Vl sublines correspond to mdr2. The mobility of the new bands appears to be identical in several different restriction digests of KB-VI and KB-Al DNA, indicating that a similar rearrangement may have occurred in both independently selected sub-15 lines. However, while the rearranged bands are amplified in KB-Al, they do not appear amplified in KB-Vl cells. In addition, both types of cells contain bands corresponding to the unrearranged allele of mdr2, which is not amplified. Amplification of the rearranged but 20 not the parental mdr2 band in KB-Al cells suggests that DNA rearrangement either preceded or occurred simultaneously with the onset of gene amplification in these In the case of KB-V1, it is unclear whether mdr2 rearrangement is related to amplification of mdrl.

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Example 6

To determine whether the evolutionarily conserved regions of mdrl and mdr2 contained transcribed sequences, pMDRl and pMDR2 were used as probes for Northern hybridization, performed according to the procedure of Thomas, Proc. Natl. Acad. Sci. (USA), 77, 5201-5205 (1980) with poly (A) RNA extracted from the parental KB-3-1 and multidrug-resistant KB-C2.5 cells [Akiyama et al., supra; Fojo et al., Cancer Res., supra; Richert et al., supra and Shen et al., supra] under the

conditions of high hybridization stringency as recited in Example 5. Poly (A) + RNA was extracted from the parental drug-sensitive KB-3-1 cells and from the colchicine-resistant KB-C2.5 subline as described in 5 Chirgwin et al., <u>Biochem.</u>, <u>18</u>, 5294 (1979). One microgram of each RNA preparation was electrophoresed in a 1.5% glyoxal agarose gel [McMaster et al., Proc. Natl. Acad. Sci. (USA), 74, 4835 (1977)] and transferred onto Gene Screen Plus™ membrane as available from New England Nuclear, Boston, MA. The membranes were hybridized with $3 \times 10^5 \text{ dpm/cm}^2$ of pMDR1 or pMDR2 probes. Hybridization was done in 1M NaCl, 10% dextran sulphate, 1% SDS, 50% formamide, 100 µg/ml denatured salmon sperm DNA at 42°C. The membranes were washed with 0.1 x SSC, 0.5% 15 SDS at 65°C and autoradiographed. The size of the RNA band was determined relative to the positions of 28S and 18S ribosomal RNA.

The probe pMDR1 hybridizes to an mRNA band of a 4.5 kb size which is highly expressed in the drugresistant cells. This mRNA is not detectable in the parental KB-3-1 cells, indicating little or no expression when the probes were labelled either by nick translation or oliogolabelling. No distinct bands, however, could be detected when pMDR2 was used as a probe. In addition, no bands were revealed by using other repeatfree subfragments of pHDR4.5 as probes in addition to pMDR2. While the existence of transcriptionally active sequences in other regions of md:2 or transcription of mdr2 sequences at a very low level cannot be excluded by these results, transcription of the amplified region of mdr2 homologous to the Chinese hamster mdr gene is not detected by Northern hybridization.

Amplification and over expression of DNA sequences homologous to the Chinese hamster mdr gene in multidrug-resistant human KB carcinoma cells suggests that a similar mechanism may be responsible for multidrug resistance in both human and rodent cells. The

nature of the proteins encoded by mdr genes is still unknown. The size of mdrl mRNA is consistent with the possibility that it may code for a 170 kd glycoprotein overexpressed in various multidrug-resistant cell lines 5 [Biedler et al., supra; Ling et al., supra; Ramu et al., supra; Beck et al., supra; Kartner et al., Science, 221, 1285 (1983); Debenham et al., Mol. Cell. Biol., 2, 881 (1982); Robertson et al., Mol. Cell. Biol., 4, 500 (1984)]. It is also unknown whether the same mechanism 10 is utilized in the development of multidrug resistance by human tumor cells in vitro and in the course of chemotherapy. The availability of cloned probes which detect transcription of mdr DNA in human cells makes it possible now to investigate expression of these sequences in clinical samples of multidrug-resistant tumors.

Example 7

- In order to examine levels of expression of mdrl sequences during the development of multidrug resistance, multidrug-resistant sublines of human KB carcinoma cells and two other human multidrug resistant cell lines of different origin were studied.
- Agents used in selecting different sublines in multiple steps were colchicine, Adriamycin and vinblastine. In the first two steps of colchicine selection, clones were only obtained if the cell populations were first mutagenized with ethylmethane sulfonate (EMS).
- 30 Similarly, KB cell lines selected independently for resistance to Adriamycin or vinblastine [Akiyama et al., supra; Fojo et al., Cancer Res., supra; Richert et al., supra; and Shen et al., supra] were obtained only after mutagenesis with EMS in the first step. Subsequent
- 35 selection, up to very high levels of resistance, was possible without mutagenesis, and occurred at high

frequency.

The isolation and some properties of the human multidrug resistant KB carcinoma cell lines has been previously described in Akiyama et al., supra; Fojo et al., Cancer Res., supra; and Richert et al., supra. The KB cell lines used in this study, the manner of their selection, and their relative resistance to various drugs, are shown in Table 3. CEM is a cell line described in Beck in Advances in Enzyme Regulation, 22, G. Weber, ed. (Pergamon Press, Oxford, 1984), 207, and 2780 is a cell line described in Rogan et al., Science, 224, 994 (1984).

To determine the extent to which mdrl sequences were expressed in these cell lines and the size of the corresponding RNAs, a Northern hybridization was performed with total RNA and poly (A)*-RNA from these cells. A 4.5 kilobase RNA, which migrates just below the 28S ribosomal RNA marker, was clearly visible in all the lanes containing either total or poly (A)*

20 RNA from the resistant lines but was not seen in any of the sensitive cell lines.

Slot blot hybridization of total RNA was used to quantitate the expression of mdrl in various sensitive and resistant cell lines. RNA prepared as previously described above was applied to filters using a Schleicher and Schuell slot blot apparatus or by blotting after electrophoresis in 1% agarose containing 13.4% formaldehyde. A gel-purified insert from the pMDRl clone was 32P-labeled for use as a probe. Nitrocellulose filters were baked and preincubated for 4-6 hours at 42°C in 50% formamide, 5 x SSC, 10X Denhardt's solution, 0.1% SDS and 100 ug/ml salmon sperm DNA. Filters were hybridized overnight in the above solution containing 32P-labeled probe. Filters were washed 3 times for 10 minutes at room temperature in 2 x SSC, 0.1% SDS and 3 times for 20 minutes at 50°C in 0.1 x

SSC, 0.1% SDS. Levels of mdrl expression were determined by densitometry of the autoradiograms. Tracings of peaks were cut out and weighed and compared to the KB-8 peak which was arbitrarily assigned a value of 1.

The results are presented in Table 3 along with the relative drug resistances of the human leukemic lymphoblast cell lines, and the human ovarian cancer cell lines used in the study. In Table 3, ND is an abbreviation for "none detected".

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TABLE

		Relative Registance	Beaista	4	- נות נות נות נות נות
Cell Line	Selecting Agent	Col	Adr	Vb1	Expression
KB-3-1	parental KB		-	7	QN
KB-8	colchicine, 5 ng/ml	2.1	1.1	1.2	1
KB-8-5	colchicine, 10 ng/ml	. ຜ ຕ	3.2	6.3	က
KB-8-5-11	colchicine, 100 ng/ml	40	23	51	80
KB-C1	colchicine, l µg/ml	260	160	96	270
KB-C1-R1	revertant of KB-Cl	9	က	4	1
KB-C1.5	colchicine, 1.5 µg/ml	320	ļ	140	340
KB-C6	colchicine, 6 µg/ml	2,100	320	370	820
KB-Al	Adriamycin, l µg/ml	19	2/5	43	270
. KB-V1	vinblastine, l µg/ml	170	4 2 0	210	320
СЕМ	parental leukemic	# T	-	г	ND
CEM-V1b ₁₀₀	vinblastine	45	120	420	250
2780	parental, ovarian	~1	~ 4	7	ND
2780-Ad	Adriamycin	1	170	15	260

As shown in Table 3, there was a good correlation between extent of multidrug resistance and the level of mdrl-specific mRNA. As can also be seen in Table 3, there is little or no expression of the mdrl sequences in parental, drug-sensitive cell lines, but increasing expression occurs as the cell lines become more resistant to drugs. A revertant cell line, KB-Cl-Rl, subcloned in the absence of colchicine from the colchicine-resistant cell line KB-Cl, still expresses mdrl sequences at reduced levels consistent with its low level of multidrug resistance.

extent of increased expression in the resistant cell lines relative to the parental line, since the hybridization signal from the parental RNA was too weak. However, the extent of expression relative to the KB-8 cell line has been calculated and these data are shown in Table 3. Expression appears to correlate well with increasing drug-resistance for every step of selection in KB cells and reaches very high levels in our most resistant KB cell lines.

The data summarized in Table 3 indicate that two other human cell lines of different origin, selected for multidrug resistance, also express high levels of the 4.5 kb mRNA. Very little or no expression of this RNA was detected in the parental cell lines. The human leukemic lymphoblast cell line CEM (A.T.C.C. CCL119) and its resistant derivatives CEM-VLB100, selected for resistance to vinblastine (gift of W. Beck, St. Jude's Hospital) (Beck, supra.) and the ovarian cell line 2780 and its resistant derivative 2780-Ad, selected for resistance to Adriamycin (gift of T. Hamilton and R. Ozols, National Institutes of Health) (Rogan et al., supra) both showed high levels of expression of the 4.5 kb mRNA. Because even low levels of cellular multidrug-resistance may result in clinically refractory tumors,

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expression of mdrl mRNA in sublines having a low level (2-6 fold) of relative drug resistance but not in the parental drug-sensitive cell lines is of particular interest. In this regard the results presented in Table 3 indicate that quantitation of mdrl mRNA expression may potentially be used for diagnosis of multidrug resistance in clinical tumor specimens.

Example 8

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To compare the levels of mdrl mRNA expression with the extent of amplification of the genomic mdrl sequences genomic DNA was isolated from all of the cell lines described in Example 6. Following digestion with HindIII, amplification of mdrl was examined by Southern blot analysis.

DNA, prepared as previously described in Example 3, was digested with HindIII and electrophoresed in 0.8% agarose gels before Southern transfer to Gene 20 Screen Plus (New England Nuclear). The blots were hybridized with the pMDRl probe for 18 hours at 42°C in 50% formamide, 5 x SSC, 1% SDS with 100 ug/ml salmon sperm DNA. The blots were then washed with 2 x SSC at room temperature for 10 minutes, 2 x SSC, 1% SDS at 42°C for 60 minutes and 0.1 x SSC at room temperature for 60 minutes prior to autoradiography.

No amplification of mdrl was found in the KB cell lines with low levels of resistance (KB-8, KB-8-5 and the revertant subline, KB-Cl-Rl), even though these cell lines expressed increased levels of mdrl mRNA. Increased expression of mdrl sequences in human cells may therefore occur prior to gene amplification. Amplification of the mdrl gene was detected in highly resistant sublines of KB cells selected in colchicine, vin-35 blastine or Adriamycin, as well as in CEM-VLB100 and 2780-Ad cell lines. In the latter two sublines, the

degree of gene amplification was estimated by densitometry to be approximately 5-10 fold for 2780-Ad and 10-15 fold for CEM-VLB $_{100}$.

In all cases, the increase in mRNA expression 5 was clearly greater than the extent of amplification. These results suggest that the evolution of these lines involved a step or steps in which expression was increased out of proportion to gene amplification. A similar dissociation of amplification and expression of the dhfr gene has been reported for human cancer cells selected for resistance to methotrexate in vitro. [Frei et-al., Proc. Natl. Acad. Sci. (USA), 81, 2875 (1984); Wolman et al., Proc. Natl. Acad. Sci. (USA), 80, 807 (1983).] The development of multidrug resistance in 15 human KB cells differs in this respect from Chinese hamster V79 cells where a low (5-7 fold) degree of relative drug resistance is accompanied by 5-10 fold amplification of mdr DNA [Roninson et al., supra; and Gros et al., supra].

20 These studies demonstrate a correlation between expression of the mdrl gene and the development of resistance to multiple agents in five independentlyderived human cell lines of different origins selected for resistance to different cytotoxic drugs. Expression 25 of mdrl may therefore represent a common mechanism of multidrug resistance in human cell lines. Increased expression of mdrl in at least some cases occurs initially without gene amplification and may be a prerequisite for the development of multidrug resistance. 30 observation may be especially relevant for the analysis of the role of the mdrl gene in the development of multidrug resistance by human tumors in the course of chemotherapy and may have diagnostic potential. Since the tumor cells are expected to have a relatively low 35 degree of resistance, such an analysis may involve quantitation of mdrl RNA expression rather than gene amplification in tumor samples.

Example 9

The segment of the mdrl gene cloned into pMDRl 5 was sequenced by the chemical degradation procedure [Maxam et al., Meth. Enzymol., 65, 499, (1980)] and the enzymatic chain-termination sequencing technique (Sanger et al., Proc. Natl. Acad. Sci. USA, 74, 5463, (1977)] using supercoiled plasmid DNA as a template [Zagursky et 10 al., Gene Anal. Techn., 2, 89, (1985)]. To facilitate sequencing, pMDRI was mapped with HaeIII and RsaI and individual 220-400 bp fragments of pMDR1 were subcloned into a pUC18 plasmid vector (Bethesda Research Laboratories, Rockville, MD). The sequence of pMDR1 was 15 confirmed by sequencing both strands. The complete sequence of pMDR1 is presented in Table 4. Comparison with the sequence of the corresponding cDNA clones in Example 10 below indicated that pMDR1 includes segments of two protein-coding sequences (exons), comprising 20 nucleotides 1-111 and 653-807, and an intervening sequence (intron) which is not expressed as mRNA and which comprises nucleotides 112-652. Table 4 shows that amino acid sequence corresponding to the exons within pMDR1. This amino acid sequence therefore defines a 25 segment of the mdrl protein product.

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TABLE 4

CLONE	
pMDR1	
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SECUENCE	

T GGA AGA CAA ATA CAC AAA ATT AGA AAA CAG TTT TTT CAT GCT Gly Arg Gln Ile His Lys Ile Arg Lys Gln Phe Phe His Ala	ATA ATG CGA CAG GAG ATA GGC TGG TTT GAT GTG CAC GAT GTT Ile MET Arg Gln Glu Ile Gly Trp Phe Asp Val His Asp Val	GGG GAG CTT AAC ACC CGA CTT ACA GA Gly Glu Leu Asn Thr Arg Leu Thr As	GTAAGTATT TAGTTTTATG TTGAACTTGG GTGTCGTTTA ATGTATTATG ATGTATTATG GTGTCGTTTA ATGTATTATG ATGTATTA GGGGGGGGATGTTA GGAGGTGTTA GTGTACTTT TATGAGACAA AATTCCTTCT AAGCAGCAAC ACTCTGCATG GCATCCTTTT GTTCCCAGTG CCTTGACAGG GTATGGGGGG ATCCTCTGAG AATGTGCATTAA ATGAAGGACT GGGCTTTCCA GAATGAAGAA TTCTGAGCAA TTTGAAATTC CTAGGTTGAA TACTTCTTTT GTACAGAAT ATGAGATATTC CTAGGTTGAA TACTTCTTTT GTACACGATG AATCTCTAG GGGCCATGT GGCCTATCGAT TACTTCTTT AATGACAAAT AAACTCTAG AAACTTCTAC CCTGCTAAAT AAAACAAAG AAACTTCTAC CCTGCTAAAT AAAACAAAAAAAAAA
1 1 0	44 4	98	112 251 251 301 4451 4451 501 601 651 74

- 25 -

ATT GGA Ile Gly GAC AAA ASP LYS ATT GGT Ile Gly Grr Val GAA Glu AAT Asn A'I'T Ile AAG Lys TCT Ser GTC Val T GAT p Asp 653

ATG TTC TTT CAG TCA ATG GCA ACA TTT TTC ACT GGG TTT ATA MET Phe Phe Gln Ser MET Ala Thr Phe Phe Thr Gly Phe Ile

969

TABLE 4 (cont'd.)

TTG	
ATT	
GTG Val	807
CTT Leu	8
ACC Thr	9
CTA	TCA Ser
AAG	CTG Leu
TGG Trp	GGA Gly
$_{\rm GGT}^{\rm GGT}$	CTT Leu
CGT Arg	GTT Val
ACA Thr	CCT Pro
TTT Phe	AGT Ser
GGA Gly	ATC Ile
GTA Val	GCC Ala
738	780

Example 10

In order to isolate cDNA clones of the mdrl gene, poly(A)+ RNA was isolated as described in Chirgwin et al., Biochemistry, 18, 5294 (1979) and Aviv et al., Proc. Natl. Acad. Sci. (USA). 69, 1408 (1972) from the subline KB-C2.5, selected with colchicine. A cDNA library was constructed using the steps of synthesizing double-stranded cDNA, blunt ending, attachment of EcoRI linkers and insertion into the phage vector Agtll [Young and Davis, supra; Huynh et al., in: DNA Cloning Techniques: A Practical Approach, D. Glover, ed., IRL Press, Oxford, (1985)]. The cDNA library was screened by plaque hybridization (Benton et al., supra) with the pMDR1 probe. Approximately 120 positive clones were isolated. The inserts from five of these clones (λ HDR5, λHDR10, λHDR28, λHDR62 and λHDR69) were re-cloned into plasmid vectors pGEM1 and pGEM4 (Promega Biotec). The partial restriction maps of these clones are shown in Fig. 3. DNA from λ HDR5 was treated with EcoRI which generated two fragments, designated 5A and 5B. fragments were subcloned into pGEM1 at its EcoRI site to give plasmids pHDR5A and pHDR5B which were deposited with the American Type Culture Collection, 12301 Parklawn Drive, Rockville, Maryland, on March 18, 1986, and which received the respective accession numbers ATCC 67040 and ATCC 67041. Similarly, $\lambda HDR10$ was treated with EcoRI and cloned into the EcoRI site of pGEMl to produce pHDR10 which was deposited with the American Type Culture Collection, 12301 Parklawn Drive, Rockville, Maryland, on March 18, 1986, as Deposit No. 67042.

To isolate the remaining portion of mdr1 cDNA, a fragment of the clone λ HDR5, indicated with a striped bar in Fig. 3, was used to screen the same cDNA library. The inserts from three of the positive clones, designated λ HDR103, λ HDR104 and λ HDR105, were re-cloned

into the EcoRI sites of plasmid vectors pGEM1 and pGEM4, giving rise to plasmids designated pHDR103, pHDR104 and pHDR105, respectively. The plasmid pHDR104 was deposited with the American Type Culture Collection, 12301 Parklawn Drive, Rockville, Maryland, on July 16, 1986, as Deposit No. 67156.

A comparison of the restriction maps of individual clones indicates divergence in the cDNA structure among, for example, clones λHDR10, λHDR28 and λHDR69. The most highly conserved region among these clones is represented by a 270 bp PvuII fragment, which corresponds to the exon regions of pMDR1 and is indicated with a solid bar above the lines in Fig. 3. The variant sequences specific to clones λHDR62 and λHDR105 were detected by DNA sequencing, and they are shown as solid bars underneath the corresponding lines in Fig. 3. In Fig. 3, sites for digestion by corresponding restriction endonucleases are identified as follows: "A", AccI; "E", EcoRI; "H", HindIII; "N", XmnI; "P", PvuII; "S", StuI; "T", SstI; "V", AvaI; and "X", XbaI.

The cDNA clones \(\text{\text{HDR10}} \), \(\text{\text{HDR5}} \) and \(\text{\text{\text{HDR104}}} \) were sequenced in their entirety using the methods of subcloning the inserts into an M13 phage vector [Messing, Meth. Enyzmol., 101, 20, 1983], generating a series of overlapping deletion subclones (Henikoff, Gene, 28, 351, 1984] and determining their DNA sequence by the enzymatic chain-termination sequencing techniques [Sanger et al., supra]. A part of the cDNA sequence was determined by specific-primer-directed DNA sequencing [Strauss et al., Anal. Biochem., 154, 353 1986] using supercoiled plasmid DNA as a template [Zagursky et al., supra]. The overlapping regions of clones λ HDR10, λ HDR5 and \alphaHDR104 were found to be identical, and therefore, these clones are assumed to represent different parts of the same cDNA. The combined cDNA sequence of clones AMDR10, AMDR5 and AMDR104 is shown in Table 5.

table also shows the amino acid sequence of $\underline{mdr}l$ gene product, derived from the same cDNA sequence.

TABLE 5

MDR1 cDNA SEQUENCE (CLONES AHDR10; AHDR5; AHDR104)

	466	508	550	592	634	919
AGATCATTTC AAAAGAGAGG GTTTCGCAGT AGTCATCTGT GGGCTGAGCA TCGAGTAGCG CCTTTAGGTC TCTTGGTGGC	AAG Lys	AAG Lys		ryr TrG Leu	CTG Leu	AAT Asn
SATCA STORY SECTO COLORY CTTT	AAG Lys	AAG Lys	CGC	ACT Thr	ATG	GGA Gly
	AAG Lys	GAT Asp	TTT	GGA Gly	ATG	GCA Ala
TAAAGATTAG GCAACCGAT AATTCAACCT GCCGGGGCGT GCCGGGCCGT CTGAGCTCAT CTGTTCGTTT	GCA Ala	AAA Lys	ATG		CTC	AAT Asn
raaag Scaac Sccc Sccc Sccc Crcct	GGA Gly	GAA Glu	TCA		CCT Pro	GCA
	GGA Gly	AGT Ser	TTT	ATG	CTT Leu	TTT Phe
TCCAGATTCC ACTTCAGGAA TCCTCCTGGA AGTCAATCCG CAGGAACAGC CACAGGAAGC GCAGAGGCCG GCAGAGGCCG	AAT Asn	AAA Lys	GTA	TAT	GGA G1y	ATC Ile
TCCA ACTI TCCI AGTC CAGG CACA GCAG	CGC Arg	AAT Asn	AGT	TTG	GCT	GAT Asp
ACTC ACTC ACTC ACTC ACTC ACTA ACTA	GAC Asp	AAC Asn	GTC	AAG	GGG G1y	ACA Thr
TCAGATATTC GCCAGAACAT ATCAGCATTC ATTGCCTGGG CTCTCTTTGC GCTCAAAGAA AGTCGGAGTA	666 61.y	CTG Leu	ACT	GAC	CAT His	ATG Met
	GAA Glu	AAA Lys	CCA	CTT	ATC Ile	GAA G1u
CCTACTCTAT TCATCCCTA TCCAACGAA TTCTCGAGGA GGTGAGGCTG CAGCGCTTCC CAGCGCTTCCAA CTTTCCACTAA	CTT Leu	TTT Phe	AAA	TGG	ATC Ile	GGA Gly
CTACT CATTC CACTC CTCC CTCTT CTCCC	GAT Asp	TTT Phe	AAG	AAT Asn	GCC Ala	TTT Phe
2422244	ATG MET	AAC Asn	GAA	TCA	GCT Ala	GTG Val
101 101 101 201 201 301 401	425	467	209	551	593	635

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<i>LL</i> 9					ATG	TCÀ	AAC	ATC	ACT	AAT	AGA	AGT	GAT	ATC	71
719	AAT Asn	GAT Asp	ACA Thr	GGG G1y	TTC Phe	TTC Phe	ASI ATG MET	AAT Asn Asn	rnr CTG Leu	Asn GAG Glu	Arg GAA Glu	Ser GAC Asp	ASP ATG MET	Ile ACC Thr	16
761	AGG Arg	TAT Tyr	GCC Ala	TAT Tyr	TAT Tyr	TAC Tyr	AGT Ser	GGA Gly	ATT Ile	GGT Gly	GCT Ala	GGG G1y	GTG	CTG	80
803	GTT Val	GCT Ala	GCT Ala	TAC Tyr	ATT Ile	CAG Gln	GTT Val	TCA	TTT Phe	TGG Trp	TGC Cys	CTG	GCA Ala	GCT Ala	84
845	GGA Gly	AGA Arg	CAA Gln	ATA Ile	CAC His	AAA Lys	ATT Ile	AGA Arg	AAA Lys	CAG Gln	TTT Phe	TTT Phe	CAT	GCT	88
887	ATA Ile	ATG MET	CGA Arg	CAG Gln	GAG Glu	ATA Ile	GGC Gly	TGG Trp	TTT Phe	GAT Asp	GTG Val	CAC His	GAT Asp	GTT Val	92
929	666 61y	GAG Glu	CTT Leu	AAC Asn	ACC Thr	CGA Arg	CTT	ACA Thr	GAT Asp	GAT Asp	GTC Val	TCT	AAĞ Lys	ATT Ile	67
971	AAT Asn	CAA Glu	GTT Val	ATT Ile	GGT Gly	GAC	AAA Lys	AT'I Ile	GGA Gly	ATG	TTC Phe	TTT	CAG Gln	TCA	101
1013	ATG MET	GCA Ala	ACA Thr	TTT Phe	rrc Phe	ACT Thr	666 G1y	TTT Phe	ATA Ile	GTA Val	GGA Gly	TTT Phe	ACA Thr	CGT Arg	105
1055	GGT Gly	TGG Trp	AAG Lys	CTA Leu	ACC Thr	CTT	GTG Val	ATT Ile	TTG	GCC Ala	ATC Ile	AGT	CCT Pro	GTT Val	109

TABLE 5 (cont'd.)

											1				
1097	CTT	GGA Gly	CTG Leu	TCA	GCT Ala	GCT Ala	GTC Val	TGG Trp	GCA Ala	AAG Lys	ATA Ile	CTA	TCT	TCA	113
1139	TTT Phe	ACT Thr	GAT	AAA Lys	GAA Glu	CTC	TTA Leu	GCG Ala	TAT Tyr	GCA Ala	AAA Lys	GCT Ala	GGA Gily	GCA	118
1181	GTA Val	GCT	GAA Glu	GAG Glu	GTC Val	TTG Leu	GCA Ala	GCA Ala	ATT Ile	AGA Arg	ACT Thr	GTG Val	ATT Ile	GCA	122
1223	TTT Phe	GCA Gly	GGA Gly	CAA Gln	AAG Lys	AAA Lys	GAA Glu	CTT Leu	GAA Glu	AGG Arg	TAC Tyr	AAC	AAA Lys	AAT Asn	126
1265	TTA Leu	GAA Glu	GAA Glu	GCT Ala	AAA Lys	AGA Arg	ATT Ile	GGG Gly	ATA Ile	AAG Lys	AAA Lys	GCT Ala	ATT Ile	ACA	130
1307	GCC Ala	AAT Asn	ATT Ile	TCT Ser	ATA Ile	GGT Gly	GCT Ala	GC'r Ala	TTC Phe	CTG Leu	CTG	ATC	TAT Tyr	GCA	134
1349	TCT Ser	TAT Tyt	GCT Ala	CTG Leu	GCC Ala	TTC Phe	TGG Trp	TAT Tyr	$\begin{array}{c} GGG\\ G1 \end{array}$	ACC Thr	ACC Thr	TTG	Grc Val	CTC	139
1391	TCA Ser	666 61y	GAA Glu	TAT Tyr	TCT Ser	ATT Ile	GGA G1y	CAA Gln	GTA Val	CTC	ACT	GTA Val	Trc Phe	TTT Phe	143
1433	TCT	GTA Val	TTA Leu	ATT Ile	GGG Gly	GCT Ala	TTT Phe	AGT Ser	GTT Val	GGA Gly	CAG Gln	GCA Ala	TCT	CCA	147
1475	AGC Se r	ATT Ile	GAA Glu	GCA Ala	TTT Phe	GCA Ala	AAT Asn	GCA Ala	AGA Arg	GGA Gly	GCA	GCT	TAT Tyr	GAA Glu	151

TABLE 5 (cont'd.)

1558	1600	1642	1684	1726	1768	1810	1852	1894	1936
\mathtt{TAT}	TTG	GAA Glu	GGG Gly	AGC	GAG Glu	AAT Asn	GAA Glu	TAT Tyr	GTC Val
AGC	AAT	AAA	AG'T	AAG	ACA	ATA	CAG	CGC	GCT
	Asin	Lys	Ser	Lys	Thr	Ile	G1n	Arg	Alà
GAC	GGA G1y	CGA Arg	CAG G1n	666 61y	CCC	ACC	AGT	ATT Ile	AAA Lys
ATT	AAG	TCT	GTG	TGT	GAC	AGG	GTG	AAC	GAG
Ile	Lys	Ser	Val	Cys	Asp	Arg	Val	Asn	Glu
AGT	ATT	CCA	AAG	GGC	TAT	ATT	GTG	GAA	ATT
Ser	Ile	Pro	Lys	Gly	Tyr	Ile	Val	Glu	Ile
CCA Pro	AAT Asn	TAC Tyr	CrG Leu	AGT Ser	CTC	GAT Asp	GGT Gly	GCT	GAG Glu
AAG	GAT	AGT	AAC	AAC	AGG	CAG	ATT	ATA	GAT
Lys	Asp	Ser	Asn	Asn		Gln	Ile	Ile	Asp
AAT	CCA	TTC	CTG	GGA	CAG	GGA	ATC	ACG	ATG
Asn	Pro	Phe	Leu	Gly	Gln	Gly	Ile		MET
GAT Asp	AAA Lys	CAC His	GGC Gly	GTT Val	ATG	GAT Asp	GAA Glu	ACC	ACC Thr
ATT	CAC	GTT	AAG	CTG	CTG	GTT	CGG	GCC	GTC
Ile	His	Val	Lys	Leu	Leu	Val	Arg		Val
ATA	666	AAT	TTG	GCC	CAG	AGT	CTA	TTT	AAT
Ile	61y	Asn		Ala	Gln	Ser	Leu	Phe	Asn
AAG	AGT	ACA	ATC	GTC	GTC	GTC	TTT	TTG	GAA
Lys	Ser	Arg	Ile	Val	Val	Val	Phe	Leu	Glu
rrc	AAG	TTC	AAG	A.C.G	ACA	ATG	AGG	GTA	CGT
Phe	Lys	Phe	Lys	Thr	Thr	MET	Arg	Val	Arg
ATC	TCG	GAA	GTT	CAG	ACA	GGG	GTA	CCT	GGC
Ile		Glu	Val	Gln	Thr	Gly	Val	Pro	Gly
1517	1559	1601	1643	1685	1727	1769	1811	1853	1895

TABLE 5 (cont'd.)

1978	2020	2062	2104	2146	2188	2230	2272	2314	2356
CAT His	AGT :	Grr ; Val	GCC ;	GAT :	CGT Arg	GAT ASP	ATG ;	ACA 7	TCC 2
Cer (Pro I	Trg 1	CTG (Leu V	TCA C	CrG (Leu A	CAT (His P	Trc C Phe P	Crc P	CAG A	GAA T
Cre	CAG Gln	GCC	ACG	GCT	GCT	GLY	GAA (ATG (GAT (ASD (
AAA	GCC	CGT	GCC	GTG	ATA	GCT	GAT	ACA	GCT
Lys	Ala	Arg	Ala	Val	Ile	Ala	Asp	Thr	
ATG	GGG	GCA	GAG	CAG	GrG	ATC	CAT	GTC	GCA
MET	Gly	Ala	Glu	Gln	Val	Ile	His	Val	Ala
ATC	AGA	ATT	GAT	GTT	ATT	GTC	AAT	CTT	AAT
Ile	Arg	Ile	Asp	Val	Ile	Val	Asn	Leu	Asn
TTT Phe	GAG Glu	GCC	CTG Leu	GTG	ACC	GAC	GGA Gly	AAA Lys	GAA Glu
GAC	GGA	ATC	CTG	GCA	ACC	GCT	AAA	TTC	TTA
	G1y	Ile	Leu	Ala	Thr	Ala	Lys	Phe	Leu
TAT	GTT	AGG	CTC	GAA	CGG	AAT	GAG	TAC	GAA
Tyr	Val	Arg		Glu	Arg	Asn	Glu	Tyr	Glu
GCC	CTG	CAG	ATC	AGC	GGT	CGT	GTG	ATT	GTT
Ala		Gln	Ile	Ser	Gly	Arg	Val	Ile	Val
AAT	ACC	AAG	AAG	GAA	AAA	GTT	ATT	GGC	GAA
Asn	Thr	Lys	Lys	Glu	Lys	Val	Tle	G1 y	Glu
GCC	GAC	CAG	CCC	ACA	AGA	ACA	стс	AAA	AAT
	Asp	Gln	Pro	Thr	Arg	Thr	Val	Lys	Asn
GAA Glu	TTT Phe	666 61y	AAC Asn	GAC Asp	GCC	TCT Ser	$\begin{array}{c} {\rm GGA} \\ {\rm G1} \end{array}$	GAC Glu	GGA Gly
AAG	AAA	GGT	CGC	TTG	AAG	\mathbf{r} rc	GAT	AAA	GCA
Lys	Lys	Gly	Arg	Leu	Lys	Leu	Asp	Lys	Ala
1937	1.979	2021	2063	21.05	2147	2189	2231	2273	2315

2398	2440	2482	2524	2566	2608	2650	2692	2734	2776
TCA	GTC Val	GAG	ATT Ile	GGT Gly	TTT Phe	ATT Ile	TCA	TTT Phe	CTC
GAT	AGT	AAA	AGG	Grr	GCA	AGA	TTT	ACA	ATC
Asp	Ser	Lys	Arg	Val	Ala	Arg	Phe	Th	Ilė
AAT Asn	AGG Arg	ACC Thr	TGG Trp	GTT Val	CCA Pro	ACA	TTG	ATT Ile	GAG G1u
TCA	CGT	AGT	TTT.	TTT	CAA	TTT	AAC	TTT	GGA
Ser	Arg	Ser	Phe	Phe		Phe	Asn	Phe	Gly
TCT Ser	ACT Thr	CTT Leu	TCC	TAT Tyr	CTG	GTT Val	AGT Ser	TCT Ser	GCT
ATG	TCA	AAG Lys	GTT Val	CCT Pro	GGC G1γ	666 61 y	AAT Asn	ATT Ile	AAA Lys
GAA	AGA	AGA	CCA	TGG	GGA	ATA	CAG	ATT	GGC
Glu		Arg	Pro	Trp	Gly	Ile	Gln	Ile	Gly
TTG	AAA	GAC Asp	CCT Pro	GAA Glu	AAT Asn	ATT Ile	CGA Arg	GGA Gly	TTT Phe
GCC	AGA	CAA	ATA	ACT	ATA	AAG	AAA	CTT	ACA
Ala	Arg	Gln	Ile	Thr	Ile	Lys	Lys	Leu	Thr
GAT Asp	ATA Ile	GCC	AGT Ser	TTA Leu	ATT Ile	TCA Ser	ACA Thr	GCC	TTC Phe
ATT	CTA	CAA	GAA	AAT	GCC	TTT	GAA	CTA	GGT
Ile	Leu	Gln	Glu	Asn	Ala	Phe	Glu	Leu	Gly
GAA	AGT	TCA	GAT	CTA	TGT	ATA	CCT	$\Upsilon \Upsilon \Upsilon$ Phe	CAG
Glu	Ser	Se r	Asp	Leu	Cys	Ile	Pro		Gln
AGT	TCC	GGA	CTG	AAG	፻ባיፓ	ATA	GAT	TTG	CTT
Ser		Gly	Leu	Lys	Phe	Ile	Asp	Leu	Leu
AAA	AGA	CGT	GCT	ATG	GTA	GCA	GAT	CTA	TTC
Lys	Arg	Arg	Ala	MET	Val	Ala	Asp	Leu	Phe
2357	2399	2441	2483	2525	2567	2609	2651	2693	2735

2818	2860	2902	2944	2986	3028	3070	3112	3154	3196
AGA Arg	GGA Gly	AAA Lys	ATA Ile	GGT Gly	ATT	CAA Gln	ATC Ile	TTG	TTG
CTC	ACT	GTT	AAT	TAT Tyr	ATC Ile	GGA G1y	AAG	TCT Ser	AGT
ATG	ACC Thr	CAA Gln	CAG Gln	ATC Ile	CCC	TCT	GGG G1y	GTT Val	¢AG Gln
TCC	AAC Asn	GCT Ala	ACC Thr	TTC Phe	GTA Val	TTG	GCT Ala	Grr Val	GCT
CGA Arg	AAA Lys	GCT Ala	ATT Ile	TCC	ATT Ile	ATG	GGT Gly	ACC	TAT Tyr
TTC Phe	CCT Pro	GAT	GTA Val	ATA Ile	GCA Ala	AAA Lys	GAA Glu	CGA Arg	ATG MET
GTT Val	GAC Asp	AAT	GCT Ala	ATT Ile	TTA Leu	ATG	CTA	TTC Phe	CAT His
ATG	GAT Asp	GCC	CTT	ATA Ile	Crc	GAA Glu	GAA Glu	AAC Asn	GAA Glu
TAC Tyr	TTT Phe	CTC Leu	AGG Arg	GGA G1y	TTA Leu	GTT Val	AAA Lys	GAA Glu	TTT Phe
CGA Arg	TGG Trp	AGG Arg	TCC	ACA Thr	CTG Leu	GTT Val	AAG Lys	ATA Ile	AAG Lys
CTC	AGT Ser	ACC Thr	GGT Gly	GGG Gly	ACA Thr	GGA G1y	GAT Asp	GCA	CAG Gln
CGG Arg	GTG Val	ACT Thr	ATA Ile	CTT Leu	CTA Leu	GCA Ala	AAA Lys	GAA Glu	GAG Glu
AAG Lys	GAT Asp	TTG Leu	GC'r Ala	AAT Asn	CAA Gln	ATA Ile	CTG	ACT Thr	CAG Gln
ACC Thr	CAG Gln	GCA Ala	GGG Gly	GCA	TGG Trp	GCA Ala	GCA Ala	GCT	ACT Thr
2777	2819	2861	2903	2945	2987	3029	3071	3113	3155

CAG Gln GGA Gly Tyr Tyr AAA Lys CGT Val	·		TAC TYr TTT Phe Cys Cys Ser GGT G1y		AAC Asn TTC Phe CGG Arg GAG Glu AET AAA	TCT Ser ACC Thr Phe GAT ASP GCC Ala			AAAA Lys ATG MET TYr TYr CAA Gln	GCA Ala ATG TTG Leu GTC Val GCA	CAC His TYR TYR Val TTT Phe Phe RGT Ser AGT	ATC Ile Phe GCA Ala TCA Ser Ser CAC	TTT Phe TCC Ser CAT His Ala TTT Phe Alc
Q 2.	ATG	ATC Ile	ATT	GAA Glu	AAA Lys	ACC Thr	CCT Pro	$\mathbf{T}\mathbf{T}\mathbf{G}$	ATT	GAC Asp		AGC	AGC TAC Ser Tyr
00	GAA Glu	66C 61y	CTA	ATG Met	CCG	AAC Asn	ACA Thr	TTC	GAA Glu	GGA Gly	A A	AAT	rr Gre
0 0	GGT Gly	GAA	GTT Val	GTA Val	TTC	AAC	TAT Tyr	CCC	ACC Thr	CGA Arg	CCG		G GAC
ບິວິວິ	GTG Val	CTT (CAG (Glu (GGA Gly	CTG	AGC Ser	CTG	GAG Glu	GTG	AAG	AAG		. GGC . G1y

3658	3700	3742	3784	3826	3868	3910	3952	3994	4036
ACA Thr	666 61y	GTT Val	CCC	GGA Gly	GCA	CCT Pro	CTC	CTT Leu	TCA
AGC	GCA Ala	AAT Asn	GAG Gl.u	TAT	AGG Arg	CTG	CAG	GCC	ACG Thu
AAG Lys	TTG Leu	CTG	CAG Gln	GCC	GTG Val	TCA	ACT Thr	CGT Arg	GCC Ala
GGG G1y	CCC	CGA Arg	TCC	ATT Ile	ATC Ile	GAG Glu	GGA Gly	GCT	GAA Glu
TGT Cys	GAC Asp	AAG Lys	GTG Val	AAC Asn	GAG Glu	ATC Ile	AAA Lys	ATA Ile	GAT Asp
GGC Gly	TAC Tyr	ATA Ile	ATC Ile	GAG Glu	GAA Glu	TTC Phe	GAC Asp	GCC	TTG
ACT Ser	TTC Phe	GAA Glu	GGC Gly	GCT Ala	CAG Gln	GCC	GGA Gly	ATT Ile	CTT
AGC	CGG Arg	AAA Lys	CTG	ATT Ile	TCA Ser	CAT His	GTA Val	CGC Arg	TTG
GGC Gly	GAG Glu	GGC Gly	CAC His	AGC	GrG Val	ATA Ile	AAA Lys	CAA Gln	ATT Ile
GTG Val	CTG Leu	GAT Asp	GCA Ala	TGC	GTG Val	AAC Asn	ACT Thr	AAA Lys	CAT' His
CTG Leu	CTC	CTT Leu	CGA Arg	GAC Asp	CGG Arg	GCC Ala	AGC Ser	CAG Gln	CCT Pro
GCT Ala	CAG Gln	CTG	CTC	TTT Phe	AGC Ser	GAG Glu	TAT Tyr	GGC Gly	CAG Gln
CTG	Grc Val	GTG Val	TGG Trp	CTG	AAC Asn	AAG Lys	AAA Lys	GGT Gly	AGA Arg
ACG Thr	GTG Val	AAA Lys	CAG Gln	ATC Ile	GAC Asp	GCA Ala	AAT Asn	TCT	Grr Val
3617	3659	3701	3743	3785	3827	3869	3911	3953	3995

4078	4120	4162	4204	4246		
CTG	CAC His	TTT Phe	CTG	CAG		TATG STAT TAAT TTTA SCCT AAAG
GCC	GCT Ala	GTG Val	CAG	Grc Vail		TTTAGATATG GAAGAGGTAT ACTTCGTAAT TCATAGTTTA TAAAAGATAA CTGACTGCCT TTGCATAAAAAAAAAA
GAA Glu	ATT Ile	GTG Val	CAG Gln	AGT		
CAA GAA Gln Glu	GTG Val	ATA Ile	CAT His	GTC Val		ATTA CAGA CAGAA GATT TGTA AATG
GTC Val	ATT Ile	TTA	ACG Th <i>r</i>	ATG MET		TTAATATTTG ACAGAATTAT GTCTTCAGAG TGGAGAGAA AAGTAGATTT TGGACTGTAA ATTGAAATGT
Grr Val	TGC	GAC	GGC Gly	TCA Ser	4267	
AAG Lys	ACC	GCA Ala	CAT His	TTT Phe	42	TAAATACTTT GCAAACACTT CAACTTCAGA CATCATCAAG AACAGAATTA TTTCCCATT AGCAAAAAGT
GAA Glu	CGC	AAT	GAG Glu	\mathtt{TAT}	TGA	TAAATACTTT GCAAACACTT CAACTTCAGA CATCATCAAG AACAGAATTA TTTCCCATT AGCAAAAAGT
AGT Ser	GGC Gly	CAG Gln	AAG Lys	ATC Ile	CAG Gln	
GAA Glu	GAA Glu	ATC Ile	GTC Val	GGC Gly	CGC Arg	TATGAGATGT AAAGTTAAAA TTFCCTCACT GAGTGAGAGA TAAATFTTAT TTTGTTATAT TTATAGAAGT
ACA Thr	AGA Arg	ACC Thr	AGA Arg	AAA Lys	AAG Lys	
GAT Asp	GCC	TCC	66C 61y	CAG Gln		ACTG ATTC AACA AACA TAAT TAAT
CTG Leu	AAA Lys	CTG Leu	AAT Asn	GCA Ala	GGA ACA Gly Thr	ACTCTGACTG ACATTTATTC CTG!!!TAACA TAAAGGAACA AAC!GCATTA AATGTGTAAT TGCTAAAAGA
GCT Ala	GAC Asp	CGC Arg	CAG Gln	CTG Leu	GCT Ala	AC TA AA AA TG
4037	4079	4121	4163	4205	4247	4268 4318 4318 4418 4468 4518 4618

Analysis of the amino acid sequence presented in Table 5 indicates that the mdrl gene product is likely to be a transmembrane protein. This protein may consist of two approximately equal parts, with a 5 considerable sequence homology to each other, indicating that the mdrl gene has likely evolved as a result of an internal duplication. Each half of the protein consists of a hydrophobic and a hydrophilic portion. Each of the hydrophobic portions includes six transmembrane domains, as determined by the algorithm of Eisenberg et al. [J. Mol. Biol., 179, 125-142 (1984)]. Both hydrophilic portions contain two regions that share a high level of amino acic homology with the ATP-binding sites of several known enzymes. The best homology has been 15 observed with the ATP-binding sites of peripheral membrane components of bacterial periplasmic binding protein-dependent transport systems [Higgins et al., EMBO J., 4, 1033-1040, (1984)]. The presence of the transmembrane domains and potential glycosylation sites 20 within the protein sequence is consistent with the mdrl protein being related to the P-glycoprotein, which is described above.

Analysis of the DNA and protein sequence information presented in Table 5 by the algorithm of 25 Eisenberg et al., supra, may be used to predict the protein regions that are located on the outside of the cell membrane. These protein regions may be produced either by chemical synthesis or by expression in the appropriate vector systems, and may be used to raise 30 antibodies against cells that express the mdrl gene product, as described in Example 11.

Example 11

well as different individual fragments of recombinant plasmids pHDR4.4 and pHDR4.5, or the latter plasmids as a whole, or cDNA clones λ HDR5, λ HDR10, λ HDR62, λ HDR28 and $\lambda HDR69$, or other sequences according to the present 5 invention, may be used as diagnostic tools for detection of human tumor cells resistant to chemotherapeutic drugs. These plasmids may be labeled directly with a radioactive isotope, according to the procedures of Rigby et al., Mol. Biol., 113, 237-251 (1977) or Feinberg et al., Anal. Biochem., 132, 6-13 (1983), for example. Alternatively, the plasmids may be labelled with a non-radioactive chemical tag, for example, according to the procedure in Leary et al., Proc. Natl. Acad. Sci. (USA), 80, 4045-4049 (1983). The plasmids may also be used to direct synthesis of labeled RNA probes [e.g., according to the procedure in Melton et al., Nucleic Acids Res., 12, 7035-7055 (1984)]. labeled probes may then be used to detect the presence of homologous RNA sequences in tumor cells either by the 20 Northern hypridization procedure (according to Thomas, Proc. Natl. Acad. Sci. (USA), 77, 5201-5205 (1980)] or by dot blot or slot blot hydridization (according to Kafatos et al., <u>Nucleic Acids Res.</u>, <u>7</u>, 1541-1552 (1979) and Brown et al., Mol. Cell. Biol., 3, 1097-1107 25 (1983)], or by in situ hybridization techniques [e.g., those according to the procedures of Brahic et al., Proc. Natl. Acad. Sci. (USA), 75, 6125-6129 (1978)]. It is anticipated that in situ hybridization will provide a particularly sensitive method for detection of a small 30 number (1 in 1000 or fewer) of multidrug-resistant cells within a biopsy.

The mdr clones may be used to obtain polyclonal or monoclonal [Yelton et al., Ann. Rev.

Biochem., 50, 657-680 (1981)] antibodies against mdr
gene products using either of two strategies.

The first strategy involves determination of

the cDNA sequences of mdr genes, as described in Example The cDNA sequence may be used to deduce the corresponding protein sequence. Peptides corresponding to different parts of mdr proteins, and preferably 5 comprising at least 15-20 amino acid residues, may be chemically synthesized by solid-phase methods [Marglin et al., Ann. Rev. Biochem., 39, 841-866 (1970)]. peptides may then be used to elicit specific polyclonal and monoclonal antibodies [Lerner, Nature, 299, 592-596 10 (1982); Niman et al., Proc. Natl. Acad. Sci. (USA), 80, 4949-4953 (1983)]. The availability of the full-length mdrl cDNA sequence, as shown in Table 5, greatly facilitates the design of potentially immunogenic peptides, corresponding to different regions of the mdrl 15 protein, including the potential extracytoplasmic domains.

The second strategy involves expression of either complete or partial mdr gene products in bacteria, yeast or mammalian expression systems using 20 plasmid, phage or viral expression vectors [Vieira et al., Gene, 19, 259-268 (1982); Young et al., Proc. Natl. Acad. Sci. (USA), 80, 1194-1198 (1983); Bitter et al., Gene, 32, 263-274 (1984); Cepko et al., Cell, 37, 1053-62 (1984); and Gorman et al., Mol. Cell. Biol. 2, 1044-25 1051 (1982)]. The expressed proteins may be purified and used in a vaccine or to raise specific antibodies. Antibodies against the mdr gene products may be used as the alternative diagnostic tools for detection of drugresistant cells. Finally, such antibodies may 30 potentially be used as a basis for a new strategy of cancer immunotherapy. This strategy may involve, for example, conjugation of anti-mdr antibodies with radioactive isotopes or chemical toxins in order to specifically eliminate multidrug-resistant tumor cells. This 35 approach may be particularly efficient if used in combination with chemotherapy. Alternatively, the binding

of anti-mdr antibodies to cells expressing mdr gene products, even in the absence of antibody-mediated cytotoxicity, may be sufficient to reverse the multidrug-resistant phenotype and may therefore render tumor cells susceptible to the cytocidal action of the chemotherapeutic drugs.

In addition, complete or partial <u>mdr</u> gene products may be used as a vaccine to elicit an immune reaction in a patient against multidrug resistant tumor cells.

Although the present invention has been described in terms of preferred embodiments, it is understood that variations and modifications will occur to those skilled in the art upon consideration of the present invention. Therefore, it is intended that all such equivalent variations and modifications should come within the scope of the invention as claimed.

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WHAT IS CLAIMED IS:

An isolated nucleic acid sequence for a
 human mdr gene associated with multidrug resistance in human cells.

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- 2. A nucleic acid selected from the group consisting of:
- (a) a nucleic acid comprising a member of the group consisting of:
- 5 a continuous sequence of nucleotides as set forth in Table 4;
 - a continuous sequence of nucleotides as set forth in Table 5;
- a continuous sequence of nucleotides as set 10 forth in pHDR4.4 (ATCC 40227);
 - a continuous sequence of nucleotides as set forth in pHDR4.5 (ATCC 40228);
 - a continuous sequence of nucleotides as set forth in pHDR5A (ATCC 67040);
- a continuous sequence of nucleotides as set forth in pHDR5B (ATCC 67041);
 - a continuous sequence of nucleotides as set forth in pHDR10 (ATCC 67042); and
- a continuous sequence of nucleotides as set 20 forth in pHDR104 (ATTC 67156);
 - (b) a nucleic acid comprising a nucleotide sequence which hybridizes with at least one of the continuous sequences of nucleotides as set forth in (a) or which is contained within the same mRNA molecule of
- 25 human origin or cDNA molecule of human origin as at least one of the continuous sequences of nucleotides as set forth in (a);
- (c) a nucleic acid comprising a nucleotide sequence which hybridizes with any nucleotide sequences30 described in (b); and
- (d) nucleic acids comprising a sequence of nucleotides which, but for the degeneracy of the genetic code, would hybridize with at least one of the continuous sequences of nucleotides as set forth in (a), (b), 35 or (c).

- 3. A nucleic acid probe comprising:
- (a) a nucleic acid comprising a member of the group consisting of:
- a continuous sequence of nucleotides as set 5 forth in Table 4;
 - a continuous sequence of nucleotides as set forth in Table 5;
 - a continuous sequence of nucleotides as set forth in pHDR4.4 (ATCC 40227);
- a continuous sequence of nucleotides as set forth in phDR4.5 (ATCC 40228);
 - a continuous sequence of nucleotides as set forth in pHDR5A (ATCC 67040);
- a continuous sequence of nucleotides as set 15 forth in pHDR5B (ATCC 67041);
 - a continuous sequence of nucleotides as set forth in pHDR10 (ATCC 67042); and
 - a continuous sequence of nucleotides as set forth in pHDR104 (ATTC 67156);
- 20 (b) a nucleic acid comprising a nucleotide sequence which hybridizes with at least one of the continuous sequences of nucleotides as set forth in (a) or which is contained within the same mRNA molecule of human origin or cDNA molecule of human origin as at
- least one of the continuous sequences of nucleotides as set forth in (a);
 - (c) a nucleic acid comprising a nucleotide sequence which hybridizes with any nucleotide sequences described in (b); and
- (d) nucleic acids comprising a sequence of nucleotides which, but for the degeneracy of the genetic code, would hybridize with at least one of the continuous sequences of nucleotides as set forth in (a), (b), or (c); and
- 35 a label associated with said polynucleotide.

or. (c).

- 4. A polypeptide comprising a continuous sequence of amino acids encoded by a nucleic acid selected from the group consisting of:
- (a) a nucleic acid comprising a member of the
 5 group consisting of:
 - a continuous sequence of nucleotides as set forth in Table 4;
 - a continuous sequence of nucleotides as set forth in Table 5;
- a continuous sequence of nucleotides as set forth in pHDR4.4 (ATCC 40227);
 - a continuous sequence of nucleotides as set forth in pHDR4.5 (ATCC 40228);
- a continuous sequence of nucleotides as set 15 forth in pHDR5A (ATCC 67040);
 - a continuous sequence of nucleotides as set forth in pHDR5B (ATCC 67041);
 - a continuous sequence of nucleotides as set forth in pHDR10 (ATCC 67042); and
- a continuous sequence of nucleotides as set forth in pHDR104 (ATTC 67156);
- (b) a nucleic acid comprising a nucleotide sequence which hybridizes with at least one of the continuous sequences of nucleotides as set forth in (a) or which is contained within the same mRNA molecule of human origin or cDNA molecule of human origin as at least one of the continuous sequences of nucleotides as set forth in (a);
- (c) nucleic acids comprising a nucleotidesequence which hybridizes with any nucleotide sequences described in (b); and
- (d) nucleic acids comprising a sequence of nucleotides which, but for the degeneracy of the genetic code, would hybridize with at least one of the continuous sequences of nucleotides as set forth in (a), (b),

5. A composition effective as a vaccine or as an antigen for induction of specific antibodies, comprising a polypeptide as recited in claim 4 and a compatible diluent, adjuvant, or carrier.

- 6. An antibody to a polypeptide comprising a continuous sequence of amino acids encoded by a nucleic acid selected from the group consisting of:
- (a) a nucleic acid comprising a member of the group consisting of:
 - a continuous sequence of nucleotides as set forth in Table 4;
 - a continuous sequence of nucleotides as set forth in Table 5;
- a continuous sequence of nucleotides as set forth in pHDR4.4 (ATCC 40227);
 - a continuous sequence of nucleotides as setforth in pHDR4.5 (ATCC 40228);
- a continuous sequence of nucleotides as set 15 forth in pHDR5A (ATCC 67040);
 - a continuous sequence of nucleotides as set forth in pHDR5B (ATCC 67041);
 - a continuous sequence of nucleotides as set forth in pHDR10 (ATCC 67042); and
- a continuous sequence of nucleotides as set forth in pHDR104 (ATTC 67156);
- (b) a nucleic acid comprising a nucleotide sequence which hybridizes with at least one of the continuous sequences of nucleotides as set forth in (a) or which is contained within the same mRNA molecule of human origin or cDNA molecule of human origin as at least one of the continuous sequences of nucleotides as set forth in (a);
- (c) a nucleic acid comprising a nucleotide 30 sequence which hybridizes with any nucleotide sequences described in (b); and
- (d) nucleic acids comprising a sequence of nucleotides which, but for the degeneracy of the genetic code, would hybridize with at least one of the continuous sequences of nucleotides as set forth in (a), (b), or (c).

- 7. A diagnostic reagent comprising:
- an antibody to a polypeptide comprising a continuous sequence of amino acids encoded by a nucleic acid selected from the group consisting of:
- 5 (a) a nucleic acid comprising a member of the group consisting of:
 - a continuous sequence of nucleotides as set forth in Table 4;
- a continuous sequence of nucleotides as set 10 forth in Table 5;
 - a continuous sequence of nucleotides as set forth in pHDR4.4 (ATCC 40227);
 - forth in pHDR4.5 (ATCC 40228);
- a continuous sequence of nucleotides as set forth in pHDR5A (ATCC 67040);
 - a continuous sequence of nucleotides as set forth in pHDR5B (ATCC 67041);
- a continuous sequence of nucleotides as set 20 forth in pHDR10 (ATCC 67042); and
 - a continuous sequence of nucleotides as set forth in pHDR104 (ATTC 67516);
- (b) a nucleic acid comprising a nucleotide sequence which hybridizes with at least one of the continuous sequences of nucleotides as set forth in (a) or which is contained within the same mRNA molecule of human origin or cDNA molecule of human origin as at least one of the continuous sequences of nucleotides as set forth in (a);
- 30 (c) a nucleic acid comprising a nucleotide sequence which hybridizes with any nucleotide sequences described in (b); and

(d) nucleic acids comprising a sequence of nucleotides which, but for the degeneracy of the genetic code, would hybridize with at least one of the continuous sequences of nucleotides as set forth in (a), (b), or (c); and

a label associated with said monoclonal antibody.

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8. An immunotherapeutic preparation comprising:

an antibody to a polypeptide comprising a continuous sequence of amino acids encoded by a nucleic 5 acid selected from the group consisting of:

- (a) a nucleic acid comprising a member of the group consisting of:
- a continuous sequence of nucleotides as set forth in Table 4;
- a continuous sequence of nucleotides as set forth in Table 5;
 - a continuous sequence of nucleotides as set forth in pHDR4.4 (ATCC 40227);
- a continuous sequence of nucleotides as set 15 forth in pHDR4.5 (ATCC 40228);
 - a continuous sequence of nucleotides as set forth in pHDR5A (ATCC 67040);
 - a continuous sequence of nucleotides as set forth in pHDR5B (ATCC 67041);
- a continuous sequence of nucleotides as set forth in pHDR10 (ATCC 67042); and
 - a continuous sequence of nucleotides as set forth in pHDR104 (ATTC 67156);
- (b) a nucleic acid comprising a nucleotide

 25 sequence which hybridizes with at least one of the continuous sequences of nucleotides as set forth in (a) or which is contained within the same mRNA molecule of human origin or cDNA molecule of human origin as at least one of the continuous sequences of nucleotides as

 30 set forth in (a);
 - (c) a nucleic acid comprising a nucleotide sequence which hybridizes with any nucleotide sequences described in (b); and

- (d) nucleic acids comprising a sequence of nucleotides which, but for the degeneracy of the genetic code, would hybridize with at least one of the continuous sequences of nucleotides as set forth in (a), (b), or (c).
 - 9. The immunotherapeutic preparation as recited in Claim 8 further comprising a cytocidal agent conjugated with said antibody.

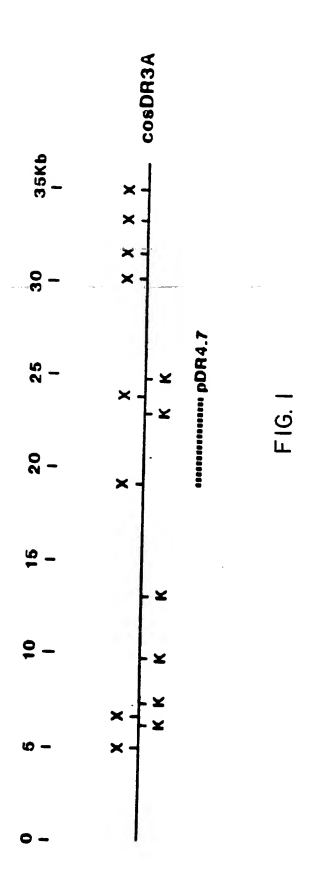
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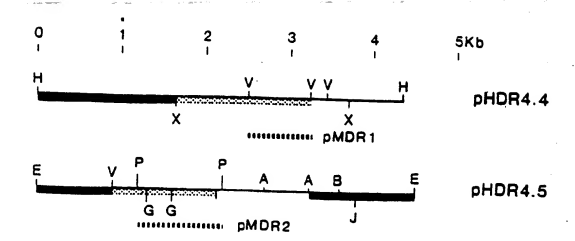
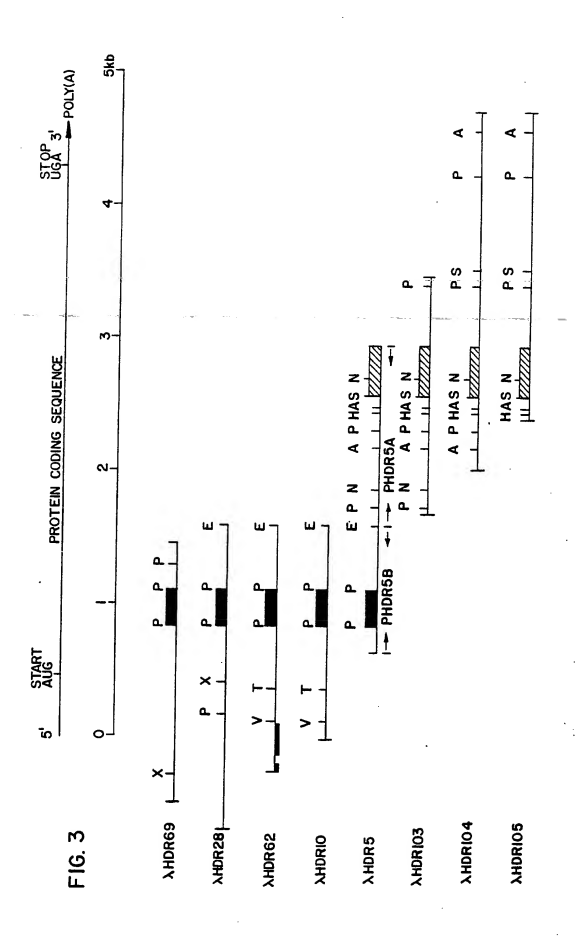


FIG. 2



SUBSTITUTE SHEET

INTERNATIONAL SEARCH REPORT

International Application No PCT/US87/00758

I. CLASSIFICATION OF SUBJECT MATTER (if several classification symbols apply, indicate all) 3								
According to International Patent Classification (IPC) or to both Nat	ional Classification and IPC							
Int. Cl. (4): Cl2Q 1/68								
U.S. C1.: 435/6								
II. FIELDS SEARCHED								
Minimum Documen	ntation Searched 4							
Classification System	Classification Symbols Classification Symbols							
435/6; 536/27								
U.S. 935/78, 9								
436/501								
Documentation Searched other	than Minimum Documentation							
	are Included in the Fields Searched 6							
Computer Search: Lexpat; APS; Biosis 1977-1987	Chemical Abstracts	1967-1987;						
III. DOCUMENTS_CONSIDERED_TO_BE_RELEVANT_14								
Category • Citation of Document, 16 with indication, where app	ropriate, of the relevant passages 17	Relevant to Claim No. 18						
Y Proceedings National Academ U.S.A., Vol. 82, No. 22, is 1985 (Washington D.C., USA) "Amplification of DNA seque multidrug-resistant KB carc pages 7661 and 7665.	sued November, A.T. FOJO ET AL. nces in human	- 1-3						
Y Proceedings National Academ U.S.A., Vol. 83 No. 2, issu (Washington D.C., USA) P. C "Isolation and characteriza sequences amplified in mult hamster cells", see page 33	ed January 1986 ROSS ET AL, tion of DNA idrug-resistant	1-3						
P,X Chemical Abstracts, Vol. 10 02 March 1987 (Columbus, OH ET AL, "The mdrl gene, resp multidrug-resistance, codes p-glycoprotein" see page 15 abstract No. 62136q, Bioche Commun. 1986, 141 (3) 956-6	onsible for for for the m. Biophys. Res.	1-3						
* Special categories of cited documents: 15 "A" document defining the general state of the art which is not considered to be of particular relevance "E" earlier document but published on or after the international filing date "L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) "O" document referring to an oral disclosure, use, exhibition or other means "P" document published prior to the international filing date but later than the priority date claimed	"T" later document published after to repriority date and not in conflicited to understand the principle invention "X" document of particular relevant cannot be considered novel or involve an inventive step "Y" document of particular relevant cannot be considered to involve document is combined with one ments, such combination being in the art. "å" document member of the same	ict with the application but e or theory underlying the ce; the claimed invention cannot be considered to ce; the claimed invention an inventive step when the or more other such docupobolious to a person skilled						
IV. CERTIFICATION								
Date of the Actual Completion of the International Search ²	Date of Mailing of this International Sc	earch Report *						
03 JUNE 1987	1 1 JUN 1987							
International Searching Authority 1	Signature of Authorized Officer 20							
TSA/IIS	Jeienz W. Jaz							

FURTHER INFORMATION CONTINUED FROM THE SECOND SHEET	į.
P,X Chemical Abstracts Vol. 105, No. 7, issued 18 August 1986 (Columbus, OH, USA) I.B. RONINSON ET AL, "Isolation of human mdr DNA sequences amplified in multidrug-resistant KB carcinoma cells" see page 165, column 2,the abstract No. 55593n, Proc. Natl. Acad. Sci. U.S.A. 1986 83(12) 4538-42 (Eng.)	
V. OBSERVATIONS WHERE CERTAIN CLAIMS WERE FOUND UNSEARCHABLE 10	
This international search report has not been established in respect of certain claims under Article 17(2) (a) for	the following reasons:
1. Claim numbers . because they relate to subject matter 12 not required to be searched by this Aut	· · · · · · · · · · · · · · · · · · ·
2. Claim numbers , because they relate to parts of the international application that do not comply w	ith the prescribed require-
ments to such an extent that no meaningful international search can be carried out 13, specifically:	
VI. NOBSERVATIONS WHERE UNITY OF INVENTION IS LACKING 11	
This International Searching Authority found multiple inventions in this international application as follows:	
i nis international Searching Authority found montple inventions in this international application as follows.	
See Attachment.	
1. As all required additional search fees were limely paid by the applicant, this international search report co	vers all searchable claims
of the international application.	vers an searchable claims
2. As only some of the required additional search fees were timely paid by the applicant, this international	search report covers only
those claims of the international application for which fees were paid, specifically claims:	
3. No required additional search fees were timely paid by the applicant. Consequently, this international sea	rch report is restricted to
the invention first mentioned in the claims; it is covered by claim numbers: 1-3.	
4. As all searchable claims could be searched without effort justifying an additional fee, the international So	earching Authority did not
invite payment of any additional fee.	·
Remark on Protest	
The additional search fees were accompanied by applicant's protest. No protest accompanied the payment of additional search fees.	

	International Application No. PCT	/US87/00758
III. DOCI	UMENTS C NSIDERED TO BE RELEVANT (CONTINUED FROM THE SEC ND SHEE	
ategory *	Citation of Document, 16 with indication, where appropriate, of the relevant passages 17	Relevant to Claim No
	Chemical Abstracts, Vol. 105, No. 3, issued 21 July 1986 (Columbus, OH, USA) J.R. RIORDAN ET AL, "Multidrug resistance in mammalian cell lines and isolation of determinant glycoprotein DNA" see page 226, column 1, the abstract No. 19855r, Eur. Pat. Appl. EP 174,810 19 MARCH 1986, GB Appl. 84/22,819 10 September 1984; 34 pp.	1-3
:	Chemical Abstracts, Vol. 101, No. 7, issued 13 August 1984 (Columbus, OH, USA) V. LING ET AL, "DNA-mediated transfer of multidrug-resistance and expression of P-glycoprotein" see page 145, column 1, the abstract No. 9.4 42p, Prog. Cancer Res. Ther. 1984 30, 53-7	1-3
	(Eng.)	<u>-</u> -
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1		

PCT/US87/00758

- VI. OBSERVATIONS WHERE UNITY OF INVENTION IS LACKING CONTINUED
- Claims 1-3, drawn to nucleic acid sequences and probes to detect a gene; class 435 subclass 6 and class 536 subclass 27.
- II. Claims 5 and 8, drawn to a vaccine and immunotherapeutic preparation; class 424 subclasses 85 and 88.
- III. Claims 4, 6 and 7, drawn to a polypeptide, an antibody against the polypeptide and a diagnostic reagent; class 436 subclass 518.

The above inventions lack unity under PCT Rule 13 since each is used for an entirely different method (i.e., hybridization, vaccination and an immunoassay).